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PATENT

MIKHAIL BAYLEY

(TYPED OR PRINTED NAME OF PERSON MAILING PAPER OR FEE)

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Attorney Docket No. SALK1650-2	PTO
CERTIFICATE OF MAILING BY "EXPRESS MAIL" "EXPRESS MAIL" MAILING LABEL NO. EL 476991607US	19/51527
DATE OF DEPOSIT February 29, 2000 I HEREBY CERTIFY THAT THIS PAPER OR FEE IS BEING DEPOSITED WITH THE UNITED STATES POSTAL SERVICE "EXPRESS MAIL POST OFFICE TO ADDRESSEE" SERVICE UNDER 37 C F.R. 1.10 ON THE DATE INDICATED ABOVE AND IS ADDRESSED TO THE ASSISTANT COMMISSIONER FOR PATENTS, WASHINGTON, D C. 20231) jez

NEW PATENT APPLICATION **CONTINUATION-IN-PART** X DIVISIONAL

ASSISTANT COMMISSIONER FOR PATENTS Box Patent Application Washington, D.C. 20231

Sir:

Transmitted herewith for filing is the divisional patent application of:

Inventors: Marc R. Montminy

METHODS FOR TREATING DIABETES MELLITUS For:

This is a request for filing a ____ continuation __X_ divisional application under 37 C.F.R. 1.53(b), of Application No. 08/961,739 filed October 31, 1997, now pending, which is a continuation-in-part of Application No. 08/194,468, filed February 10, 1994, issued on May 12, 1998, as U.S. Patent No. 5,750,336.

FULL NAME OF FIRST INVENTOR	LAST NAME:	FIRST NAME:	MIDDLE NAME:			
	Montminy	Marc	R.			
CITIZENSHIP	STATE OR FOREIGN COUNTRY: United States					
POST OFFICE ADDRESS	POST OFFICE ADDRESS: 1002 Quail Garden Court	CITY AND STATE: Encinitas, California	<i>ZIP CODE</i> : 92024			

The issue fee has been paid in the above-identified application, however, it is not yet issued.

- Cancel in this application original claims 8-11 and 13-16, of the prior application 1. before calculating the filing fee. (At least one original independent claim must be retained for filing purposes.)
- 2. A preliminary amendment is enclosed.

Marc R. Montminy

Application No.: Unassigned Filed: February 29, 2000

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PATENT Attorney Docket No.: SALK1650-2

The filing fee has been calculated as shown below:

For	Number Filed		Number Extra		Rate			F	'ee
					Small Entity	Other Entity		Small Entity	Other Entity
Total Claims	9 -20	11	0	х	\$9	\$18	11	\$.00	\$.00
Independent Claims	3 -3	=	0	Х	\$39	\$78	#	\$.00	\$.00
Multiple Dependent Claims Presented: Yes	_X_No				\$130	\$260		\$00	\$.00
			BASIC	FEE	\$690	\$760		\$690.00	\$
				то	TAL FEE		\$690.00	\$	

- 3. X The Assistant Commissioner is hereby authorized to charge a total payment of \$690.00 for the filing fee, and any other fees associated with this communication or credit any overpayment to Deposit Account No. 07-1895. A duplicate copy of this Transmittal Sheet is enclosed.
 - X Any additional filing fees required under 37 C.F.R. 1.16.
 - X Any patent application processing fees under 37 C.F.R. 1.17.
- 4. X Amend the specification by inserting before the first paragraph on page 1:

This application is a ____ continuation _X_ divisional of Application No. 08/961,739 filed October 31, 1997, now pending, which is a continuation-in-part of Application No. 08/194,468, filed February 10, 1994, issued on May 12, 1998, as U.S. Patent No. 5,750,336, the entire contents of which are hereby incorporated by reference herein.

- 5. X A verified statement claiming small entity status was filed in parent application No. 08/194,468, filed February 10, 1994, and such status is still proper.
- 6. X The prior application is assigned of record to The Salk Institute for Biological Studies.
- 7. X The power of attorney in the prior application is to Stephen E. Reiter, Registration No. 31,192.

Marc R. Montminy

Application No.: Unassigned Filed: February 29, 2000

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8. <u>X</u> Please transfer the drawings from the prior application to the new application.

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Attorney Docket No.: SALK1650-2

- 9. <u>X</u> Information Disclosure Statements filed in the prior application under 37 C.F.R. 1.97 are hereby made of record (copies of 1449's and 892's are enclosed herewith for the Examiner's convenience).
- 10. \mathbf{X} Please transfer the computer readable form (CRF) copy of the Sequence Listing from the prior application, which CRF copy was filed with a Communication mailed October 5, 1999, to this new application.
- Please transfer the Statement under 37 C.F.R. § 1.821(f) and (g) from the prior 11. Xapplication, which Statement was filed with a Communication mailed October 5, 1999, to this new application.
- A true copy of the prior application as filed is enclosed, including the 12. <u>X</u> Declaration and Power of Attorney filed in parent application, U.S. Serial No. 08/194,468, filed February 10, 1994.

Address all future communications to:

Stephen E. Reiter GRAY CARY WARE & FREIDENRICH LLP 4365 Executive Drive, Suite 1600 San Diego, California 92121-2189 Telephone: (858)-677-1409

Facsimile: (858)-677-1465

The undersigned states that the enclosed application papers comprise a copy of the prior application as filed.

Respectfully submitted,

Date: <u>February 29, 2000</u>

Stephen E. Reiter Attorney for Applicant Registration No. 31,192 Phone: (858) 677-1409

Fax: (858) 677-1465

GRAY CARY WARE & FREIDENRICH LLP 4365 Executive Drive, Suite 1600 San Diego, CA 92121-2189 Gray Cary\GT\6165294.1

62574-990000

Attorney Docket No.: SALK1651 Applicant or Patentee: Marc R. Montminy Serial No. or Patent No.: 08/961,739

Filed: October 31, 1997

METHODS FOR TREATING DIABETES MELLITUS

VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 C.F.R. SS1.9(f) and 1.27(d) - NONPROFIT ORGANIZATION

I hereby declare that I am an official empowered to act on behalf of the nonprofit organization identified below:

NAME OF ORGANIZATION ADDRESS OF ORGANIZATION

THE SALK INSTITUTE FOR BIOLOGICAL STUDIES

10010 NORTH TORREY PINES ROAD
LA JOLLA CALJEORNIA 92037

LA JOLLA, CALIFORNIA 92037
TYPE OF ORGANIZATION University or other Institution of Higher Education X Tax Exempt under Internal Revenue Service Code (26 U.S.C. §§501(a) and 501(c) (3)) Nonprofit Scientific or Educational under Statute of State of the United States of America (Name of State) (Citation of Statute) Would qualify as tax exempt under Internal Revenue Service Code (26 U.S.C. §§501(a) and 501(c) (3)) if located in the United States of America
Would qualify as nonprofit Scientific or Educational under Statute of State of the United States of America if located in the United States of America (Name of State) (Citation of Statute)
I hereby declare that the nonprofit organization identified above qualifies as a nonprofit organization as defined in 37 C.F.R. §1.9(e) for purposes of paying reduced fees under Section 41(a) and (b) of Title 35, United States Code, with regard to the invention entitled <u>METHODS FOR TREATING DIABETES MELLITUS</u> by inventor(s Marc R. Montminy described in:
□ the specification filed herewith X application Serial No. 08/961,739, filed October 31, 1997 □ Patent No, issued
I hereby declare that rights under contract or law have been conveyed to and remain with the nonprofit organization with regard to the above-identified invention.
If the rights held by the nonprofit organization are not exclusive, each individual, concern or organization having rights to the invention is listed below and no rights to
the invention are held by any person, other than the inventor, who could not qualify as a small business concern under 37 C.F.R. §1.9(d) or by any concern which would not
qualify as a small business concern under 37 C.F.R. §1.9(d) or a nonprofit organization under 37 C.F.R. §1.9(e).
NOTE: Separate verified statements are required from each named person, concern or organization having rights to the invention averring to their status as small
entities (37 C.F.R. §1.27).
Full Name
Address Individual Small Business Concern Nonprofit Organization
Full Name
Address Individual Small Business Concern Nonprofit Organization
☐ Individual ☐ Small Business Concern ☐ Nonprofit Organization
Full Name
Address Individual Small Business Concern Nonprofit Organization
· ·
I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate (37 C.F.R. §1.28(b)).
I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and
further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section
1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to
which this verified statement is directed.
NAME OF PERSON SIGNING Douglas D. Busch
TITLE IN ORGANIZATION Assistant Secretary & Director of Legal Services and Technology Transfer
ADDRESS OF PERSON SIGNING 10010 North Torrey Pines Road, La Jolla, CA 92037
SIGNATURE DATE Jamay 27, 1998
GT\6055908.1 62574.991651

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:) Group Art Unit: Unassigned Examiner: Unassigned Marc R. Montminy CERTIFICATE OF MAILING BY "EXPRESS MAIL" Application No.: Unassigned) "EXPRESS MAIL" MAILING LABEL NO.) EL 476991607US DATE OF DEPOSIT FEBRUARY 29, 2000, I HEREBY CERTIFY THAT Filed: February 29, 2000 THIS PAPER OR FEE IS BEING DEPOSITED WITH THE UNITED STATES POSTAL SERVICE "EXPRESS MAIL POST OFFICE TO ADDRESSEE" SERVICE UNDER 37 C.F.R. 1.10 For: METHODS FOR TREATING ON THE DATE INDICATED ABOVE AND IS ADDRESSED TO THE ASSISTANT COMMISSIONER FOR PATENTS, **DIABETES MELLITUS** WASHINGTON, D.C. 20231 MIKHAIL BAYLEY (SIGNATURE OF PERSON MAILING PAPER OF FEEY

Box Patent Application Assistant Commissioner of Patents Washington, D.C. 20231

PRELIMINARY AMENDMENT

Sir:

This Preliminary Amendment is being filed prior to examination of the aboveidentified application. This Amendment accompanies a request under 37 C.F.R. § 1.53(b) to file a divisional application based on Application No. 08/961,739, filed October 31, 1997, now pending.

Marc R. Montminy

Application No.: Unassigned Filed: February 29, 2000

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PATENT Docket No.: SALK1650-2

IN THE CLAIMS

Please amend claims 12 and 17 as noted below. For the Examiner's convenience, all pending claims are presented, with those not being amended at this time marked "reiterated."

- 1. (Reiterated) A method for treating diabetes mellitus, said method comprising contacting a biological system with an effective amount of a compound which inhibits binding of CREB to CBP.
- 2. (Reiterated) A method according to claim 1 wherein said treatment of diabetes mellitus ameliorates hyperglycemia.
 - 3. (Reiterated) A method according to claim 2 wherein gluconeogenesis is modulated.
- 4. (Reiterated) A method according to claim 3 wherein transcription of PEPCK is inhibited.
- 5. (Reiterated) A method according to claim 2 wherein transcription of glucogon gene is inhibited.
- 6. (Reiterated) A method according to claim 1 wherein said biological system is an intact organism.
- 7. (Reiterated) A method according to claim 1 wherein said contacting is carried out by oral, intravenous, subcutaneous, intramuscular or intracutaneous mode of administration.

Marc R. Montminy

Application No.: Unassigned Filed: February 29, 2000

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PATENT Docket No.: SALK1650-2

- (Amended) A method for treating diabetes mellitus, comprising contacting a 12. biological system with an effective amount of a compound [identified by the method of claim 8] which disrupts complex comprising cyclic AMP response element binding protein (CREB) and CREB binding protein (CBP), said compound identified by a method comprising:
 - contacting a modified host cell with a test compound, wherein said modified host cell comprises:

a first fusion protein comprising a GAL4 DNA binding domain. operatively associated with the kinase-inducible domain (KID) of CREB,

a second fusion protein comprising an activation domain, operatively associated with the CREB binding domain (KIX) of CBP, and

a reporter construct comprising a GAL4 response element operatively linked to a reporter gene; and

- selecting those test compounds which cause reduced expression of the <u>(b)</u> reporter gene product, wherein said compounds are identified as disrupting complex comprising CREB and CBP.
- 17. (Amended) A method for treating diabetes mellitus, comprising contacting a biological system with an effective amount of a compound [identified by the method of claim 13] which disrupts complex comprising cyclic AMP response element binding protein (CREB) and CREB binding protein (CBP), said compound identified by a method comprising:
 - contacting a modified host cell with a test compound, wherein said modified host cell comprises:

a first fusion protein comprising an activation domain, operatively associated with the kinase-inducible domain (KID) of CREB.

a second fusion protein comprising a GAL4 DNA binding domain, operatively associated with the CREB binding domain (KIX) of CBP, and

Marc R. Montminy

Application No.: Unassigned

Filed: February 29, 2000

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a reporter construct comprising a GAL4 response element operatively

PATENT

Docket No.: SALK1650-2

linked to a reporter gene; and

(b) selecting those test compounds which cause reduced expression of the reporter gene product, wherein said compounds are identified as disrupting complex comprising CREB and CBP.

REMARKS

By the present communication, claims 12 and 17 have been amended to define Applicant's invention with greater particularity. No new matter is added by the subject amendment as all amended claim language is fully supported by the specification and original claims. Accordingly, claims 1-7, 12 and 17 are pending.

It is believed that the application is in condition for allowance and, therefore, prompt and favorable action is earnestly solicited. If there are any questions concerning this communication, the Examiner is invited to call the undersigned at the telephone number provided below.

Respectfully submitted,

Date: <u>February 29, 2000</u>

Stephen E. Retter Reg. No. 31,192

Telephone: (858) 677-1409 Facsimile: (858) 677-1465

GRAY CARY WARE & FREIDENRICH LLP 4365 Executive Drive, Suite 1600 San Diego, California 92121-2189

Gray Cary\ GT\6129662.2 62574-990000 CERTIFICATE OF MAILING BY "EXPRESS MAIL"

"EXPRESS MAIL LABEL NUMBER EM299800764US DATE OF DEPOSIT October 31, 1997

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TAE KIM
(TYPED OR PRINTED NAME OF PERSON MAILING PAPER)

(SIGNATURE OF PERSON MAILING PAPER OR FEE)

APPLICATION

for

UNITED STATES LETTERS PATENT

on

METHODS FOR TREATING DIABETES MELLITUS

by

Marc R. Montminy

Number of Drawings: Two

Docket No.: SALK 1651 Salk File No.: S97037

Attorneys

Gray Cary Ware & Freidenrich 4365 Executive Drive, Suite 1600 San Diego, California 92121-2189

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METHODS FOR TREATING DIABETES MELLITUS

RELATED APPLICATIONS

This application is a continuation-in-part of United States Serial No. 08/194,468, filed April 11, 1994, now pending, incorporated by reference herein in its entirety.

ACKNOWLEDGMENT

This invention was made in part with Government support under Grant No. GM 37828 provided by the National The Government may have certain Institutes of Health. rights in this invention.

FIELD OF THE INVENTION

The present invention relates to analytical In a particular aspect, the present invention methods. relates to methods for the identification of compounds 15 which mediate the interaction between signal dependent transcription factors and co-factor protein(s) involved in the activation of transcription. In another aspect, the present invention relates to methods for the identification of new signal dependent transcription factors. another aspect, the present invention relates to methods for the identification of novel co-factor protein(s) which dependent interaction between signal mediate the transcription factors and inducer molecules involved in the activation of transcription. In yet another aspect, the present invention relates to methods for treating diabetes mellitus.

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BACKGROUND OF THE INVENTION

Many eukaryotic genes are requlated inducible, cell type-specific fashion. Genes expressed in response to heat shock, steroid/thyroid hormones, phorbol esters, cyclic adenosine monophosphate (cAMP), factors and heavy metal ions are examples of this class. The activity of cells is controlled by external signals that stimulate or inhibit intracellular events. process by which an external signal is transmitted into and within a cell to elicit an intracellular response is referred to as signal transduction. Signal transduction is generally initiated by the interaction of extracellular factors (or inducer molecules, i.e., growth factors, hormones, adhesion molecules, neurotransmitters, and other mitogens) with receptors at the cell surface. Extracellular signals are transduced to the inner face of the cell membrane, where the cytoplasmic domains receptor molecules contact intracellular targets. initial receptor-target interactions stimulate a cascade of additional molecular interactions involving multiple disseminate intracellular pathways that the signal throughout the cell.

Many of the proteins involved in signal transduction contain multiple domains. Some of these domains have enzymatic activity and some of these domains are capable of binding to other cellular proteins, DNA regulatory elements, calcium, nucleotides, lipid mediators, and the like.

Protein-protein interactions are involved in all stages of the intracellular signal transduction process - at the cell membrane, where the signal is initiated in the cytoplasm by receptor recruitment of other cellular proteins, in the cytoplasm where the signals are disseminated to different cellular locations, and in the

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nucleus where proteins involved in transcriptional control congregate to turn on or turn off gene expression.

Mitogenic signaling affects the transcriptional activation of specific sets of genes and the inactivation of others. The nuclear effectors of gene activation are transcription factors that bind to DNA as homomeric or heteromeric dimers. Phosphorylation also modulates the function of transcription factors, as well. Oncogenes, first identified as the acute transforming genes transduced by retroviruses, are a group of dominantly acting genes. Such genes, which are involved in cell division, encode growth factors and their receptors, as well as second messengers and mitogenic nuclear proteins activated by growth factors.

The binding of growth factors to their respective receptors activates a cascade of intracellular pathways phospholipid requlate metabolism, arachidonate metabolism, protein phosphorylation, calcium mobilization and transport, and transcriptional regulation. Specific phosphorylation events mediated by protein kinases and phosphatases modulate the activity of a variety transcription factors within the cell. These signaling events can induce changes in cell shape, mobility, and adhesiveness, or stimulate DNA synthesis. Aberrations in these signal-induced events are associated with a variety of hyperproliferative diseases ranging from cancer to psoriasis.

The ability to repress intracellular signal-induced response pathways is an important mechanism in negative control of gene expression. Selective disruption of such pathways would allow the development of therapeutic agents capable of treating a variety of disease states related to improper activation and/or expression of specific transcription factors. For example, in patients

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with non-insulin dependent diabetes mellitus (NIDDM), hyperglycemia develops, in part as a result of β cell failure secondary to chronic insulin resistance. This hyperglycemia appears to be exacerbated by hyperglucogonemia and increased hepatic gluconeogenesis. cAMP appears to be the major starvation state signal which triggers glucagon gene expression as well as transcription of PEPCK, the rate limiting enzyme in gluconeogenesis.

There remains, thus, a need in the art for 10 selective disruption of intracellular signal-induced response pathways.

SUMMARY OF THE INVENTION

In accordance with the present invention, it has been discovered that CREB binding protein (CBP) cooperates with upstream activators involved in the activation of transcription by signal dependent transcription factors, such as c-Jun (responsive to phorbol ester), serum response factor, and the like. Accordingly, assays employing CBP have been developed for the identification of compounds which disrupt the ability of signal dependent transcription factors to activate transcription. In another aspect, CBP have been developed assays employing signal dependent transcription identification of new In yet another aspect of the present invention, factors. have been developed employing CBP assays identification of novel co-factor protein(s) which mediate the interaction between signal dependent transcription factors and inducer molecules involved in the activation of In still another aspect, an assay is transcription. provided to identify compounds which have the binding and/or activation properties characteristic of CREB binding In still another aspect, methods employing protein. compounds which inhibit intracellular signal-induced

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response pathways have been developed for the treatment of diabetes mellitus.

BRIEF DESCRIPTION OF THE FIGURES

summarizing bar graph Figure 1 is a injections described in Example 2. Each bar represents the percentage of positive cells expressing β -galactosidase from 2-3 experiments where 100-200 cells were injected in [anti-CBP] denotes concentration of each experiment. affinity purified CBP antiserum injected into cells. (hatched bars) indicate the percent lacZ positive cells 10 microinjection CRE-lacZ reporter with after of antiserum (anti-CBP) or control IgG (RbIgG). Preincubation of antisera with CBP peptide or non-specific ILS peptide (1mg/ml) was carried out as indicated.

Figure 2 is a bar graph summarizing the results of CBP antisera injections, as described in Example 3. Bars represent the percentage of lacZ positive (blue) cells (mean \pm standard deviation) from 3-5 experiments where 100-Injected cells 200 cells were injected in each experiment. 20 were identified by immunofluorescence and/or lacZ staining. encoding the lacZ reporter plasmid Reporter microinjected into NIH3T3 cells. CRE-, SRE-, reporter activities were determined after microinjected cells were treated as described herein. CMV-, RSV-, and SV40-lacZ reporter activities were measured in the absence Hatched bars indicate % blue cells after of inducers. microinjection with CBP antiserum. Solid bars indicate % blue cells following injection with control rabbit IgG (RbIgG).

DETAILED DESCRIPTION OF THE INVENTION

Cyclic AMP (cAMP) regulates the transcription of numerous genes through protein kinase-A (PK-A) mediated

phosphorylation, at Ser133, of transcription factor CREB. Within the CREB protein, a 60 amino acid Kinase Inducible Domain (KID) mediates transcriptional induction by PK-A. Based on recent work describing a nuclear CREB Binding it has been examined whether CBP 5 Protein (CBP), necessary for cAMP regulated transcription. Within CBP, a CREB binding domain has been identified, referred to as KIX which specifically interacts with the phosphorylated KID domain of CREB. About 600A of solvent accessible surface area in each protein is directly involved in formation of 10 CREB:CBP complex. Phosphorylated Ser133 coordinates with a single arginine residue (Arg-600). The apparent Kd of the CREB:CBP complex is 0.4 μ M.

found to CBP have been against Antisera completely inhibit transcription from a cAMP responsive 15 promoter, but not from constitutively active promoters. Surprisingly, CBP has also been found to cooperate with upstream activators involved in phorbol ester and serum It is demonstrated herein that responsive transcription. recruitment of CBP to certain inducible promoters is intimately involved in transmitting inductive signals from phosphorylated, and thus activated, upstream factors to the RNA polymerase II complex. A number of analytical uses for CBP and CBP-like compounds based on these observations are described herein.

In accordance with the present invention, there is provided a method for the identification of a compound which inhibits activation of cAMP and mitogen responsive genes, said method comprising:

monitoring expression of reporter in response to exposure to said compound, relative to expression of reporter in the absence of said compound,

> wherein exposure to said compound is carried out in the presence of:

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- a signal dependent transcription factor,
- a polypeptide comprising at least amino acid residues 461-661 of the protein set forth in SEQ ID NO:2, and
- a reporter construct comprising a reporter gene under the control of said signal dependent transcription factor.

As employed herein, the phrase "cAMP and mitogen responsive genes" refers to early response genes which are activated in response to a diverse array of growth factors, mitogens, such as, including differentiation inducers and biomodulators. Examples of such agents include insulin-like growth factor (IGF-1), erythropoietin (EPO), nerve growth factor (NGF), epidermal growth factor (EGF), platelet-derived growth factor (PDGF), fibroblast growth factor (FGF), transforming growth factor factor tumor necrosis (TNF), interferon, (TGFS), granulocyte-macrophage colony-stimulating interleukins, factor (GM-CSF), G-CSF, prolactin, serotonin, angiotensin, noradrenalin, putrescine, bradykinin, bombesin, concanavalin A, various oncogenic agents including tumor estrogen, progesterone, irradiation, UV viruses, testosterone, glucagon, PEPCK and the like.

Signal dependent transcription factors contemplated for use in the practice of the present invention include phosphorylation dependent activators such as CREB, Jun, Fos, and other early response genes such as Myc, Myb, erbA, and Rel, serum responsive factor, Elk, as well as steroid hormone receptors (e.g., glucocorticoid receptor (GR)), and the like.

Polypeptides employed in the invention assay function as co-factors by binding to the signal dependent transcription factor as a necessary component transcriptionally active complex. Examples of such co-factors include CBP (i.e., substantially the entire amino acid sequence set forth in SEQ ID NO:2),polypeptide comprising amino acid residues 1-661 as set forth in SEQ ID NO:2, as well as functional fragments thereof, e.g., residues 461-661, and homologues thereof, such as those identified by the method described herein for the identification of compounds which have the binding and/or activation properties characteristic of CREB binding In accordance with one embodiment of the present protein. there are provided purified and invention, polypeptides, CBPs, that bind to a specific sequence within phosphorylated CREB.

As used herein, the term "purified" means that the molecule is substantially free of contaminants normally associated with a native or natural environment. CREB binding protein, or functional fragments thereof, useful in the practice of the present invention, can be obtained by a number of methods, e.g., precipitation, gel filtration, ion-exchange, reversed-phase, DNA affinity chromatography, and the like. Other well-known methods are described in Deutscher et al., Guide to Protein Purification: Methods in Enzymology Vol. 182, (Academic Press, 1990), which is incorporated herein by reference.

Alternatively, a purified CBP, or functional fragment thereof, useful in the practice of the present invention, can also be obtained by well-known recombinant methods as described, for example, in Ausubel et al., Current Protocols in Molecular Biology (Greene Publishing Associates, Inc. and John Wiley & Sons, Inc. 1993), also incorporated herein by reference. An example of recombinant means to prepare CBP, or functional fragments

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thereof, is to express nucleic acid encoding CBP, functional fragment thereof, in a suitable host cell, such as a bacterial, yeast or mammalian cell, using methods well known in the art, and recovering the expressed protein, again using methods well known in the art.

CBPs, and biologically active fragments thereof, useful in the practice of the present invention can also be produced by chemical synthesis. Synthetic polypeptides can be produced using Applied Biosystems, Inc. Model 430A or 431A automatic polypeptide synthesizer and chemistry provided by the manufacturer. CBP, and biologically active fragments thereof, can also be isolated directly from cells which have been transformed with the expression vectors described below in more detail.

The present invention also encompasses nucleic acids encoding CBP and functional fragments thereof. for example, SEQ ID NO:1. This invention also encompasses nucleic acids which encode substantially the entire amino acid sequence set forth in SEQ ID NO:2 (for example, the 20 nucleic acid sequence set forth in SEQ ID NO:1, as well as nucleic acid sequences which differ from that set forth in SEQ ID NO:1 due to the degeneracy of the genetic code), nucleic acids which encode amino acid residues 1-661, as set forth in SEQ ID NO:2, nucleic acids which encode amino acid residues 461-661, as set forth in SEQ ID NO:2, as well as nucleic acids which encode substantially the same amino acid sequences as any of those referred to above, but which differ only by the presence of conservative amino acid changes that do not alter the binding and/or activation 30 properties of the CBP or CBP-like polypeptide encoded thereby.

The invention further provides the abovedescribed nucleic acids operatively linked to a promoter, as well as other regulatory sequences. As used herein, the term "operatively linked" means positioned in such a manner that the promoter will direct the transcription of RNA from the nucleic acid. Examples of such promoters are SP6, T4 and T7.

Vectors which contain both a promoter and a 5 cloning site into which a piece of DNA can be inserted so as to be operatively linked to the promoter are well known Preferably, these vectors are capable of in the art. transcribing RNA in vitro or in vivo. Examples of such vectors are the pGEM series (Promega Biotech, Madison, WI). 10 This invention also provides a vector comprising a nucleic acid molecule such as DNA, cDNA or RNA encoding a CBP polypeptide. Examples of additional vectors useful herein are viruses, such as bacteriophages, baculoviruses and retroviruses, cosmids, plasmids, and the like. Nucleic 15 acids are inserted into vector genomes by methods well For example, insert and vector DNA can known in the art. restriction enzyme to a exposed to complementary ends on both molecules that base pair with each other and which are then joined together with a 20 ligase. Alternatively, synthetic nucleic acid linkers that correspond to a restriction site in the vector DNA can be ligated to the insert DNA which is then digested with a restriction enzyme that recognizes a particular nucleotide Additionally, an oligonucleotide containing a sequence. 25 termination codon and an appropriate restriction site can be ligated for insertion into a vector containing, for example, some or all of the following: a selectable marker gene, such as neomycin gene for selection of stable or cells; transient transfectants in mammalian 30 enhancer/promoter sequences from the immediate early gene of transcription; for high levels human CMV transcription termination and RNA processing signals from polyoma origins mRNA stability; SV40 replication and ColE1 for proper episomal replication; versatile multiple cloning sites; and T7 and SP6 RNA

promoters for *in vitro* transcription of sense and antisense RNA. Other means are available and can readily be accessed by those of skill in the art.

Also provided are expression vectors comprising encoding a mammalian CBP, or functional fragment thereof, adapted for expression in a bacterial cell, a yeast cell, a mammalian cell or other animal cell. vectors comprise the regulatory elements necessary for expression of the DNA in the bacterial, yeast, mammalian or Regulatory elements are positioned relative animal cells. 10 to the DNA encoding the CBP polypeptide so as to permit Regulatory elements required for expression thereof. to bind RNA sequences promoter include expression polymerase and transcription initiation sequences for For example, a bacterial expression ribosome binding. 15 vector includes a promoter such as the lac promoter and the Shine-Dalgarno sequence and the start codon AUG (Ausubel et al., supra 1993) for transcription initiation. Similarly a eukaryotic expression vector includes a heterologous or homologous promoter for RNA polymerase II, a downstream 20 the start codon AUG, polyadenylation signal, termination codon for detachment of the ribosome. vectors can readily be obtained commercially or assembled by methods well known in the art, for example, the methods described above for constructing vectors in general. 25 Expression vectors are useful to produce cells that express CBP or functional fragments thereof.

As employed herein, the term "reporter construct" refers to a recombinant construct, for example, an expression vector comprising a reporter gene under the control of a signal dependent transcription factor. In yet another example, the term refers to an expression vector comprising a reporter gene under the control of GAL4 response element. A compound which induces activation or inactivation of a target gene induces the reporter gene to

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express an exogenous identifiable "signal". Expression of the reporter gene indicates that the target gene has been modulated. Exemplary reporter genes encode luciferase, β -galactosidase, chloramphenicol transferase, and the like.

5 In practicing the assays of the invention, reporter plasmid is introduced into suitable host cells, along with CBP or a CBP-like polypeptide (or a construct encoding same) and signal dependent transcription factor. The transfected host cells are then cultured in the presence and absence (as a control) of test 10 compound suspected of being capable of inhibiting activation of cAMP and mitogen responsive genes. Next the transfected and cultured host cells are monitored for induction (i.e., the presence) of the product of the 15 reporter gene.

In accordance with the present invention, expression of the reporter gene can be monitored in a variety of ways. Immunological procedures useful for in vitro detection of a polypeptide in a sample include immunoassays that employ a detectable antibody. Such immunoassays include, for example, ELISA, microfluorimetric assay, agglutination assays, cytometry, serum diagnostic assays and immunohistochemical staining procedures which are well known in the art. antibody can be made detectable by various means well known in the art. For example, a detectable marker can be directly or indirectly attached to the antibody. include, for example, radionuclides, fluorogens, chromogens and chemiluminescent labels.

In accordance with still another embodiment of the present invention, there are provided methods to identify compounds which inhibit activation of cAMP and mitogen responsive genes, preferably compounds which

disrupt complex comprising CREB and CBP, said method comprising:

(a) contacting a modified host cell with a test compound, wherein said modified host cell comprises:

a first fusion protein comprising a GAL4 DNA binding domain, operatively associated with the KID domain of CREB,

a second fusion protein comprising an activation domain, operatively associated with the KIX domain of CBP, and

a reporter construct comprising a GAL4 response element operatively linked to a reporter gene; and

(b) selecting those test compounds which cause reduced expression of the reporter gene product.

In a preferred embodiment of the present invention, the first fusion protein comprises a GAL4 DNA binding domain, operatively associated with CREB and/or the second fusion protein comprises an activation domain operatively associated with CBP.

As used herein, the term "disrupt" embraces compounds which cause substantially complete dissociation of the various components of the complex, as well as compounds which merely alter the conformation of one or more components of the complex so as to reduce the repression otherwise caused thereby.

Any cell line can be used as a suitable "host" for the functional bioassay contemplated for use in the practice of the present invention. Thus, cells contemplated for use in the practice of the present

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invention include transformed cells, non-transformed cells, neoplastic cells, primary cultures of different cell types, and the like. Exemplary cells which can be employed in the practice of the present invention include Schneider cells, CV-1 cells, HuTu80 cells, F9 cells, NTERA2 cells, NB4 cells, HL-60 cells, 293 cells, Hela cells, yeast cells, NIH3T3 cells and the like. Preferred host cells for use in the functional bioassay system are COS cells and CV-1 cells. COS-1 (referred to as COS) cells are monkey kidney cells that express SV40 T antigen (Tag); while CV-1 cells 10 do not express SV40 Tag. The presence of Tag in the COS-1 derivative lines allows the introduced expression plasmid to replicate and provides a relative increase in the amount of receptor produced during the assay period. are presently preferred because they are particularly 15 convenient for gene transfer studies and provide a sensitive and well-described host cell system.

The above-described cells (or fractions thereof) are maintained under physiological conditions when contacted with physiologically active compound. "Physiological conditions" are readily understood by those of skill in the art to comprise an isotonic, aqueous nutrient medium at a temperature of about 37°C.

Various constructs employed in the practice of the present invention are well known in the art. Thus, the GAL4 DNA binding domain, the activation domain and GAL4 response elements have all been well characterized and extensively discussed in the art. For example, the DNA binding domain of the yeast GAL4 protein comprises at least the first 74 amino acids thereof (see, for example, Keegan et al., Science 231:699-704 (1986)). Preferably, the first 90 or more amino acids of the GAL4 protein will be used, with the first 147 amino acid residues of yeast GAL4 being presently most preferred.

Activation domains contemplated for use in the practice of the present invention are well known in the art and can readily be identified by the artisan. Examples include the GAL4 activation domain, BP64, VP16, and the like.

Exemplary GAL4 response elements are those containing the palindromic 17-mer:

5'-CGGAGGACTGTCCTCCG-3' (SEQ ID NO:4),

such as, for example, 17MX, as described by Webster et al.,
in Cell 52:169-178 (1988), as well as derivatives thereof.
Additional examples of suitable response elements include
those described by Hollenberg and Evans in Cell 55:899-906
(1988); or Webster et al. in Cell 54:199-207 (1988).

"operatively used herein. the phrase As associated with" means that the respective DNA sequences 15 (represented, for example, by the terms "GAL4 response element" and "reporter gene") are operational, i.e., work for their intended purposes; the word "functionally" means that after the two segments are linked, upon appropriate activation by a ligand-receptor complex, the reporter gene 20 will be expressed as the result of the fact that the corresponding "response element" was "turned otherwise activated.

As readily recognized by those of skill in the 25 art, the above-described assay can be modified to facilitate identification of compounds which inhibit any of the specific interactions involved in the formation of the CREB:CBP complex.

Compounds which are capable of inhibiting 30 activation of cAMP and mitogen responsive genes, and hence can be identified by the invention assay method, include

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antibodies raised against the binding domain of the protein set forth in SEQ ID NO:2, antibodies raised against the binding domain of CBP-like compounds, and the like. Presently preferred antibodies are those raised against a polypeptide fragment comprising amino acid residues from about 461 up to 661 of the protein set forth in SEQ ID NO:2; with antibodies raised against a polypeptide fragment comprising amino acid residues from about 634 up to 648 of the protein set forth in SEQ ID NO:2 (this subfragment is also set forth specifically as SEQ ID NO:3), especially preferred. Alternatively, antibodies which are raised against the amino acid residues surrounding residue 600 of CBP (see SEQ ID NO:2) or antibodies which inhibit the phosphorylation of residue 133 of CREB are also desired (see, for example, Parker et al., Mol Cell Biol (1996) 16(2):694-703).

Antibodies contemplated for use in the practice of the present invention have specific reactivity with the above-described CBP or CBP-like compounds. Active antibody encompassed within the definition fragments are "antibody." As used herein "specific reactivity" refers to the ability of an antibody to recognize and bind to an epitope on CBP or CBP-like compounds. Antibodies employed in the practice of the present invention can be produced by any method known in the art. For example, polyclonal and monoclonal antibodies can be produced by methods well known in the art, as described, for example, in Harlow and Lane, A Laboratory Manual (Cold Spring Harbor Antibodies: incorporated Laboratory 1988), which is The above-described CBP or CBP-like compounds reference. can be used as the immunogen in generating such antibodies. Altered antibodies, such as chimeric, humanized, CDRgrafted or bifunctional antibodies can also be produced by methods well known to those skilled in the art. antibodies can also be produced by hybridoma, chemical or recombinant methodology described, for example in Ausubel

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The antibodies can be used for determining et al., supra. a CBP-derived polypeptide, for the presence of purification of CBP-derived polypeptides, for in vitro diagnostic methods, and the like.

Alternative compounds which are capable inhibiting activation of cAMP and mitogen responsive genes polypeptide fragments comprising amino include residues from about 461 up to 661 of the protein set forth in SEQ ID NO:2. Polypeptide fragments comprising amino acid residues set forth specifically as SEQ ID NO:3 or KIX polypeptide fragments having a mutation at residue 600 (Arg-600), set forth in SEQ ID NO:2, are preferred, while KIX polypeptide fragments substituting Gln for Arg-600 are presently most preferred. Other polypeptide fragments contemplated for use in the practice of the invention include those comprising the KID domain, preferably those comprising residue 133 of CREB. In the most preferred CREB polypeptide fragment, serine residue 133 is mutated to an amino acid residue which can not be 20 phosphorylated. Other compounds which inhibit activity (i.e., phosphorylated-Ser133) by binding to CBP include adenovirus E1A oncoprotein (Nakajima et al. Genes Dev (1997) 11(6):738-747), and the like. Those of skill in the art will readily recognize other polypeptide fragments which will readily inhibit the formation of CREB: CBP complex employing such assays as those disclosed herein.

In accordance with another embodiment of present invention, there is provided a method for the identification of a compound which inhibits activation of cAMP and mitogen responsive genes, said method comprising:

> (1) contacting a test system with said compound under physiological conditions; and

(2)	monitoring expression of reporter	in
	response to said compound, relative	to
	expression of reporter in the absence	of
	said compound, wherein said reporter	is
	encoded by a reporter construct comprise	ing
	a reporter gene under the control of	a
	signal dependent transcription factor, a	and

wherein said test system comprises:

said signal dependent transcription
factor,

a polypeptide comprising at least amino acid residues 461-661 of the protein set forth in SEQ ID NO:2, and

said reporter construct.

In accordance with yet another embodiment of the present invention, there is provided a method for the identification of a compound which promotes activation of cAMP and mitogen responsive genes, said method comprising:

monitoring expression of reporter in response to exposure to said compound, relative to expression of reporter in the absence of said compound,

wherein exposure to said compound is carried out in the presence of:

- a signal dependent transcription factor, or
- a polypeptide comprising at least amino acid residues 461-661 of the protein set forth in SEQ ID NO:2, and

a reporter construct;

wherein said reporter construct comprises a reporter gene under the control of a signal dependent transcription factor.

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In accordance with still another embodiment of the present invention, there is provided a method for the identification of a compound which has the binding and/or activation properties characteristic of CREB binding protein, said method comprising:

monitoring expression of reporter in response to exposure to said compound, relative to expression of reporter in the absence of said compound,

wherein exposure to said compound is carried out in the presence of:

> signal dependent transcription factor, and

a reporter construct,

wherein said reporter construct comprises a reporter gene under the control of a signal dependent transcription factor.

In accordance with a still further embodiment of the present invention, there is provided methods for the identification of a compound which has the transcription 20 activation properties characteristic of a signal dependent transcription factor, said method comprising:

monitoring expression of reporter in response to exposure to said compound, relative to expression of reporter in the absence of said compound,

> wherein exposure to said compound is carried out in the presence of:

> > polypeptide comprising at least amino acid residues 461-661 of the protein set forth in SEQ ID NO:2, and

a reporter construct,

wherein said reporter construct comprises a reporter gene under the control of a signal dependent transcription factor.

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In accordance with a still further embodiment of the present invention, there are provided methods for treating diabetes mellitus, said method comprising contacting a biological system with an amount of 5 effective amount of a compound which inhibits binding of CREB to CBP. Such methods ameliorate hyperglycemia modulating associated with diabetes mellitus by gluconeogenesis and/or hyperglucagonemia. Particularly, such methods employ compounds which disrupt the formation of CREB:CBP complexes, thus inhibiting transcription of PEPCK or glucogon gene.

employed herein, the phrase "biological system" refers to an intact organism or a cell-based system containing the various components required for response to 15 the ligands described herein, e.g., an isoform of RAR (i.e., RAR α , RAR β or RAR γ), a silent partner for the RAR isoform (e.q., RXR), and an RAR-responsive reporter (which typically comprises an RAR response element (RARE) operative communication with a reporter gene; suitable 20 reporters include luciferase, chloramphenicol transferase, β -galactosidase, and the like.

Contacting in a biological system contemplated by the present invention can be accomplished in a variety of ways, and the treating agents contemplated for use herein can be administered in a variety of forms combination with a pharmaceutically acceptable carrier therefor) and by a variety of modes of delivery. Exemplary pharmaceutically acceptable carriers include carriers oral, suitable for intravenous, subcutaneous. 30 intramuscular, intracutaneous, and the like administration. Administration in the form of creams, lotions, tablets, dispersible powders, granules, syrups, elixirs, sterile aqueous or non-aqueous solutions, suspensions or emulsions, and the like, is contemplated.

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For the preparation of oral liquids, suitable carriers include emulsions, solutions, suspensions, syrups, and the like, optionally containing additives such as wetting agents, emulsifying and suspending sweetening, flavoring and perfuming agents, and the like.

For the preparation of fluids for parenteral administration, suitable carriers include sterile aqueous non-aqueous solutions, suspensions, or emulsions. Examples of non-aqueous solvents or vehicles are propylene glycol, polyethylene glycol, vegetable oils, such as olive oil and corn oil, gelatin, and injectable organic esters Such dosage forms may also contain such as ethyl oleate. adjuvants such as preserving, wetting, emulsifying, dispersing agents. They may be sterilized, for example, by 15 filtration through a bacteria-retaining incorporating sterilizing agents into the compositions, by compositions, the or by heating irradiating compositions. They can also be manufactured in the form of sterile water, or some other sterile injectable medium immediately before use.

As employed herein, the phrase "effective amount" refers to levels of compound sufficient to circulating concentrations high enough to modulate the expression of gene(s) mediated by members 25 steroid/thyroid superfamily of receptors. Such concentration typically falls in the range of about 10 nM up to 2 μ M; with concentrations in the range of about 100 nM up to 500 nM being preferred. Since the activity of different compounds described herein may vary considerably, and since individual subjects may present a wide variation in severity of symptoms, it is up to the practitioner to determine a subject's response to treatment and vary the dosages accordingly.

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The invention will now be described in greater detail by reference to the following non-limiting examples.

EXAMPLE I Functional Properties of CBP

To characterize the functional properties of CBP, rabbit CBP antiserum was developed against a fragment of CBP extending from amino acid residues 634-648 within the CREB binding domain of CBP (i.e., KVEGDMYESANSRDE; SEQ ID Crude antiserum was affinity purified on a NO:3). synthetic CBP peptide column, as described by Gonzalez et al., in Mol. and Cell Biol. 11(3):1306-1312 (1991), which is incorporated herein by reference. Far-Western and Western blot assays were performed as described by, for example, Chrivia et al., in Nature 365:855-859 (1993), also incorporated herein by reference. Thus, Western (CBP) and Far-Western (32P-CREB) blot analysis of partially purified CBP protein from HeLa nuclear extract was carried out following SDS-PAGE and transfer to nitrocellulose. Western blots were also obtained for crude HeLa nuclear 20 extracts using 32P-labeled CREB, phosphorylated with PK-A or casein kinase II (CKII). Far-Western blot analysis was also conducted with immunoprecipitates prepared from HeLa nuclear extracts with control IgG or affinity purified CBP antiserum (CBP-Ab). CREB binding activity was detected with ³²P-labeled CREB phosphorylated with PK-A.

Using the above-described antiserum, a 265 kD polypeptide was detected on Western blots, as predicted from the cDNA (see Chrivia et al., supra), which coincided with the predominant phospho-CREB binding activity in HeLa nuclear extracts by "Far-Western" blot assay. An identical phospho-CREB binding activity was also found in NIH3T3 This phospho-CREB binding protein appeared to be specific for Ser133 phosphorylated CREB because no such band was detected with CREB labeled to the same specific

activity at a non-regulatory phospho-acceptor site (Ser156) by casein kinase II (CKII) (see Hagiwara et al., Cell 70:105-113 (1992),which is incorporated herein by reference).

To further demonstrate that the major phospho-CREB binding protein in HeLa and NIH3T3 cells anti-CBP specifically bound by the antibody, immunoprecipitates were prepared from crude nuclear extracts using the CBP antiserum. Far-Western analysis of 10 these immunoprecipitates revealed a 265 kD band in samples incubated with CBP antiserum, but not with control IgG.

EXAMPLE II Role of Phosphorylation in CREB-CBP Interaction

To examine whether the phosphorylation dependent 15 interaction between CREB and CBP was critical for cAMP responsive transcription, a microinjection assay was employed using CBP antiserum, which would be predicted to impair formation of a CREB-CBP complex. Thus, NIH3T3 cells were cultured in 5% CO2 atmosphere in Dulbecco's Modified 20 Eagle's Medium (DMEM), supplemented with 10% fetal calf Forty-eight hours prior to injection, cells were passaged into scored glass coverslips and made quiescent by incubation in medium containing 0.05% fetal calf serum for 24 hours (see, for example, Hagikara et al., supra and Alberts et al., in Mol. and Cell Biol. 13:2104-2112 (1993), both incorporated herein by reference). Representative fields of NIH3T3 cells were injected with pCRE-lacZ reporter plasmid plus 5, 0.5, and 0.05 mg/ml of affinity purified CBP antiserum. Total antibody concentration in 30 microinjected cells was maintained at 5 mg/ml by adjusting with control Rabbit IgG. Injected cells were stimulated with 0.5 mM 8-bromo-cAMP, plus 3-isobutyl-1-methylxanthine (IBMX) for 4 hours, then fixed and assayed for lacZ

activity (β -Gal) as well as antibody content (Texas Red anti-Rb).

Following microinjection into nuclei of NIH3T3 cells, a CRE-lacZ reporter was markedly induced by treatment with 8-bromo-cAMP plus IBMX. Co-injection of CBP antiserum with the CRE-lacZ plasmid inhibited cAMP dependent activity in a dosage-dependent manner, but control IgG had no effect on this response.

To determine whether CBP antiserum inhibited cAMP 10 responsive transcription by binding specifically to CBP, peptide blocking experiments were performed. Thus, the effect of CBP antiserum on CRE-lacZ reporter activity following pre-treatment of CBP antiserum with synthetic CBP peptide (anti-CBP+CBP) or unrelated peptide (anti-CBP+ILS; the unrelated peptide, ILS, is described by Leonard et al., 15 in Mol. Endocr. 7: 1275-1283 (1993), which is incorporated herein by reference) was determined. Rabbit IgG+CBP and rabbit IgG pre-treated with CBP peptide were used as NIH3T3 cells were injected with CRE-lacZ controls. reporter plus various CBP antisera, stimulated with 0.5 mM 20 8-bromo-cAMP, plus IBMX for 4 hours, and assayed for lacZ Cells expressing the lacZ gene product form a activity. blue precipitate upon X-gal staining, which quenches immunofluorescent detection of the injected antibody.

25 CBP antiserum, pre-incubated with synthetic CBP peptide, was unable to recognize the 265 kD CBP product on a Western blot, and could not inhibit CRE-lacZ reporter activity upon microinjection into NIH3T3 cells. But antiserum treated with an unrelated synthetic peptide (ILS) retained full activity in both Western and microinjection assay, suggesting that the ability of the antiserum to bind CBP was critical for its inhibitory effect on cAMP dependent transcription.

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Results of these experiments are summarized in Figure 1.

EXAMPLE III Multiple Signaling Pathways Utilize CBP

determine whether CBP 5 TO activity restricted to a subset of promoters, several constitutively active reporter constructs were tested:

> Cytomegalovirus (CMV-lacZ), Rous sarcoma virus (RSV-lacZ), and SV40 (SV40-lacZ).

Thus, cells were microinjected with CBP antiserum plus Rous Sarcoma Virus (pRSV-lacZ) or Cytomegalovirus (pCMV-lacZ) reporter constructs. Alternatively, NIH3T3 microinjected with CBP antiserum (or non-specific rabbit 15 IgG (RbIgG)), plus reporter constructs containing either cAMP responsive elements (pCRE-lacZ), serum responsive elements (pSRE-lacZ) or phorbol ester responsive elements Light field photo-micrographs show cells (pTRE-lacZ). stained for β -galactosidase activity following four hour 20 treatment with either 0.5 mM 8-bromo-cAMP, plus IBMX (pCRElacZ), 20% fetal calf serum (pSRE-lacZ), or 200ng/ml TPA (pTRE-lacZ). Results of β-qalactosidase assays summarized in Figure 2. Dark field photos microinjected IgGs as visualized by immunofluorescence using Texas Red donkey anti-rabbit IgG.

When examined in NIH3T3 cells by transient transfection assay, each of the constitutively active reporter constructs had comparable basal activity, relative the cAMP-stimulated CRE reporter plasmid, thereby permitting the effects of CBP antiserum on these reporters compared directly. Although co-injected CBP antiserum could block cAMP stimulated activity from a CRElacZ reporter in contemporaneous assays, no inhibition was observed on basal expression from any of the constitutive

promoter constructs tested, even when 10-fold lower amounts of reporter plasmid were employed.

These results suggest that CBP can indeed discriminate between basal and signal dependent activities in vivo.

EXAMPLE IV

CBP-involvement in non-CREB mediated pathways

Previous reports showing that serum and phorbol stimulate their esters target genes through 10 phosphorylation-dependent trans-activators (see, for example, Hill et al., in Cell 73:395-406 (1993) or Smeal et al., in Nature 354:494-496 (1991), both incorporated herein by reference), suggested that CBP might also function in these signaling pathways. Thus, Far-Western analyses were 15 carried out with crude HeLa nuclear extracts using 32Plabeled recombinant Jun protein phosphorylated in vitro with either Jun-kinase (JNK; see Hibi et al., in Genes and Develop. 7:2135-2148 (1993), incorporated herein reference) or casein kinase II (CK II).

Whereas serum and TPA could stimulate reporter activity in NIH3T3 cells microinjected with serum responsive element (SRE)-lacZ and TPA-responsive element (TRE)-lacZ indicator plasmids, respectively, co-injected CBP antiserum completely blocked both responses. These results suggest that CBP not only interacts with CREB, but also with other signal-dependent transcription factors.

In this regard, phorbol esters and serum induce TRE-dependent transcription, in part, through the Junkinase (JNK) mediated phosphorylation of c-Jun at Ser63 and Ser73 (see, for example, Smeal et al., supra or Hibi et al., supra). Using ³²P-labeled recombinant c-Jun protein,

phosphorylated at Ser63 and Ser73 with JNK, Far-Western blot assays were performed on crude HeLa nuclear extracts. JNK-phosphorylated c-Jun protein could bind CBP with comparable affinity to CREB. But c-Jun labeled to similar 5 specific activity at non-activating sites (Thr 231, Ser243, and Ser249; see Boyle et al., in Cell 64:573-584 (1991)) by could not interact with CBP, suggesting that interaction between CBP and c-Jun requires phosphorylation of the transcriptionally active Ser63 and Ser73 phosphoacceptor sites. In view of the inhibitory effect of CBP antiserum on TRE- β gal reporter expression following phorbol ester and serum induction, the phosphorylation dependent interaction between CBP and c-Jun would appear to be a critical component of these response pathways.

EXAMPLE V Chromatographic purification of CBP

the surprising discovery that Based on cooperates with phosphorylation dependent activators by recruiting general transcription factors to promoters, it was next examined whether CBP would cofractionate with any general factors in HeLa nuclear Thus, Far-Western analyses of protein fractions obtained after phospho-cellulose chromatography. Phospho-CREB binding proteins visualized were ³²P-labeled CREB phosphorylated in vitro with PK-A Western analysis was carried out with the same $(^{32}P-CREB)$. blot as described above, using affinity purified CBP Far-Western (32P-CREB) and Western antibody (CBP Ab). (CBP-Ab) analyses of fractions were also carried out following DEAE and DE52 chromatography. Phosphocellulose, DEAE, and DE52 chromatography was performed on HeLa nuclear extracts as described by Ferreri et al., in Proc. Natl. Acad. Sci. USA in press (1993), which is incorporated herein by reference.

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Both CBP-immunoreactive and phospho-CREB binding activities were retained on phosphocellulose columns and Further purification of a were eluted at 0.3-0.5M KCl. comparable phospho-cellulose fraction on DEAE-sepharose and resins showed that CBP was highly enriched fractions containing (E, F, H) but not Although the general factor which associates activities. directly with CBP is not known, the co-fractionation of CBP with proteins involved in basal transcription initiation suggests a testable mechanism for CBP action. the results presented herein suggest that particular, phosphorylation-dependent activators like CREB and Jun influence assembly of late-acting factors (TFII E, F, transcriptional initiation/reinitiation by during interacting with CBP in a signal dependent manner.

While the invention has been described in detail with reference to certain preferred embodiments thereof, it will be understood that modifications and variations are within the spirit and scope of that which is described and claimed.

SEQUENCE LISTING

SEQ ID NO:1

[[The full nucleic acid and amino acid sequence of CBP to be provided here...]]

SEQ ID NO:2 Deduced amino acid sequence of CBP

SEQ ID NO:3 KVEGDMYESANSRDE

SEQ ID NO:4

5'-CGGAGGACTG TCCTCCG-3'

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A A C # L 7 & C R E H R H V L H & H 1144 1464 1513 1444 141 CA CTA AND THE ATT WA MA TO ACT CTT CTA ACT TO AND THE ATT OF A ACT CTT CTA ACT TO 1726 L T # P L H # b . . . 1434 53 P T A A P P 3 8 T 6 V A A 6 M R E 8 V T Q 3 L A 3 L 412 Y A I A S S Y E G B B I E A A S A A F 2432 9 F C L E E A A A T A L E X 9 6 1 L E X 9 7 A L AGSTAFFEGF SIAL MAT FCA TITE AME CCA AFR FCC CTO DOA AME OFF CAM TITE CCA CAM DCA CCC AFR DEA CC 2240 1374 B331 CTB CCA CM6 AMC CM6 TTT CCA TCA TCC AGT GOG OCA ATB AGT GTB AMC AGT BTB GOC ATG GOG CAA CCA GCA GCC CM6 GCA GGT GTT TCA CMG GOT CAG GAA CCT GOA 163 L P G H G Q P A A Q A G V B Q G Q L P G ... A CCC CAS SOC AGE CAS CTS CCT TOC CCA CCA STS ACA CAS TCA CCA TTS CAC COS ACT CCA CCT CCT CCT TCC ACA ... 1 0CA CA4 0CT ... TA CHE CAN CAA CCA ACH CCT GTG CAT ACT E A E F 4 6 Q P 4 P P V 3624 LENKT 3451 CM4 TAT STG SAT SAT SAT ATC ANG CTT ATS TTC ANC ANT SOS TOG CTA TAT MAT CST ANA ACS TCC CST STA TAT ANA TIT TO 1161 G T V B B V R L H F R H A H L T H B R T H A V I H F C 3412

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That which is claimed is:

- 1. A method for treating diabetes mellitus, said method comprising contacting a biological system with an effective amount of a compound which inhibits binding of CREB to CBP.
- 2. A method according to claim 1 wherein said treatment of diabetes mellitus ameliorates hyperglycemia.
 - 3. A method according to claim 2 wherein gluconeogenesis is modulated.
- A method according to claim 3 wherein
 transcription of PEPCK is inhibited.
 - 5. A method according to claim 2 wherein transcription of glucogon gene is inhibited.
 - 6. A method according to claim 1 wherein said biological system is an intact organism.
- 7. A method according to claim 1 wherein said contacting is carried out by oral, intravenous, subcutaneous, intramuscular or intracutaneous mode of administration.
- 8. A method for identification of a compound 20 which inhibits activation of cAMP and mitogen responsive genes, said method comprising:
 - (a) contacting a modified host cell with a test compound, wherein said modified host cell comprises:
 - a first fusion protein comprising a GAL4 DNA binding domain, operatively associated with the KID domain of CREB,

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a second fusion protein comprising an activation domain, operatively associated with the KIX domain of CBP, and

a reporter construct comprising a GAL4 response element operatively linked to a reporter gene; and

- (b) selecting those test compounds which cause reduced expression of the reporter gene product.
- 9. A method according to claim 8, wherein said GAL4 DNA binding domain is operatively associated with CREB.
- 10. A method according to claim 8, wherein said activation domain is operatively associated with CBP.
- 11. A method according to claim 8 wherein compounds which disrupt complex comprising CREB and CBP are identified.
- 12. A method for treating diabetes mellitus, comprising contacting a biological system with an effective amount of a compound identified by the method of claim 8.
- 13. A method to identify compounds which disrupt complex comprising CREB and CBP, said method comprising:
 - (a) contacting a modified host cell with a test compound, wherein said modified host cell comprises:
 - a first fusion protein comprising an activation domain, operatively associated with the KID domain of CREB,

10	a second fusion protein comprising a
	GAL4 DNA binding domain, operatively
	associated with the KIX domain of CBP, and
	a reporter construct comprising a GAL4
	response element operatively linked to a
15	reporter gene; and

- (b) selecting those test compounds which cause reduced expression of the reporter gene product.
- 14. A method according to claim 13, wherein said 20 activation domain is operatively associated with CBP.
 - 15. A method according to claim 13, wherein said GAL4 DNA binding domain is operatively associated with CBP.
- 16. A method according to claim 13 wherein 25 compounds which disrupt complex comprising CREB and CBP are identified.
 - 17. A method for treating diabetes mellitus, comprising contacting a biological system with an effective amount of a compound identified by the method of claim 13.

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ABSTRACT

METHODS FOR TREATING DIABETES MELLITUS

In accordance with the present invention, it has been discovered that CREB binding protein (CBP) cooperates with upstream activators involved in the activation of transcription of such signal dependent transcription factors as c-Jun (responsive to phorbol ester), response factor, and the like. It has also been discovered that CBP can be employed in an assay to identify compounds which disrupt the ability of such signal transcription factors to activate transcription. another aspect, it has been discovered that CBP can be employed in an assay to identify new signal dependent In yet another aspect of the transcription factors. present invention, it has been discovered that CBP can be employed in an assay to identify novel co-factor protein(s) 15 which mediate the interaction between signal dependent transcription factors and inducer molecules involved in the activation of transcription. Accordingly, the present invention provides methods for the identification of compounds which inhibit activation of cAMP and mitogen 20 responsive genes and methods for the identification of novel signal dependent transcription factors and co-factor proteins. In still another aspect, methods employing compounds which inhibit intracellular signal-induced response pathways have been developed for the treatment of 25 diabetes mellitus.

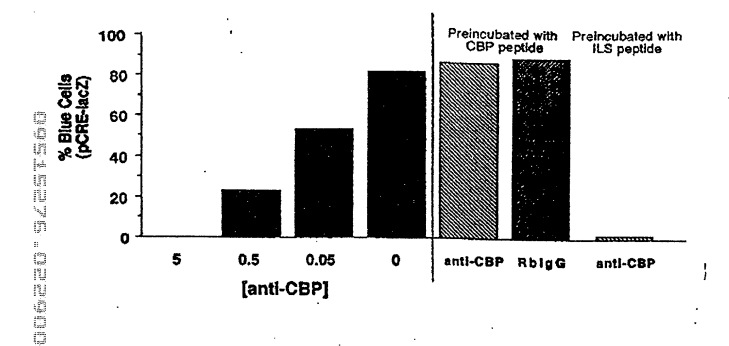


FIGURE 1

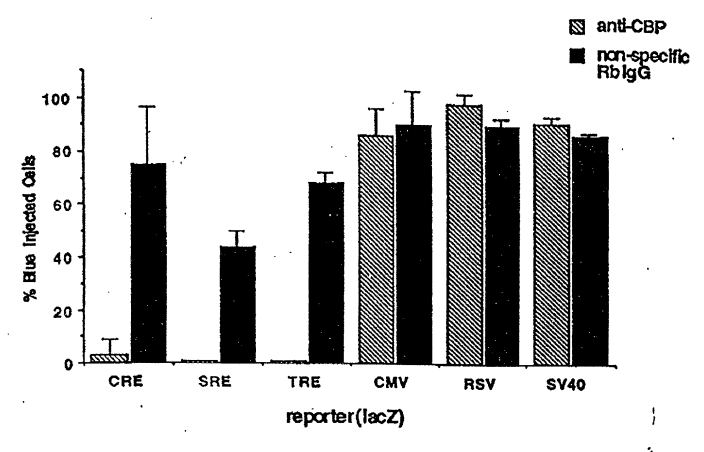


FIGURE 2

DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

As the below-named inventors, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled METHODS FOR TREATING DIABETES MELLITUS the specification of which

_____ is attached hereto.

_____ was filed on October 31, 1997 (Attorney
Docket No. SALK1651) as Application Serial
No. 08/961,739.

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment(s) referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, Sec. 1.56(a).

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

I hereby appoint the following attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith:

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POWER OF ATTORNEY BY ASSIGNEE

As a below-named assignee of the above-identified application ("Application"):

I hereby appoint the following attorneys of the assignee to prosecute the Application and to transact all business in the United States Patent and Trademark Office connected therewith:

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In re Application of:
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Application No.

Application No.: Filed: Herewith

Page 2

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Date:		

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